

1    **Transposon mutagenesis in *Pseudomonas fluorescens* reveals genes involved in blue pigment**  
2    **production and antioxidant protection.**

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15    Declarations of interest: none

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## 17    **Abstract**

18    *Pseudomonas fluorescens* Ps\_77 is a blue-pigmenting strain able to cause food product discoloration,  
19    causing relevant economic losses especially in the dairy industry. Unlike non-pigmenting *P.*  
20    *fluorescens*, blue pigmenting strains previously were shown to carry a genomic region that includes  
21    homologs of *trpABCD* genes, pointing at a possible role of the tryptophan biosynthetic pathway in  
22    production of the pigment. Here, we employ random mutagenesis to first identify the genes involved  
23    in blue-pigment production in *P. fluorescens* Ps\_77 and second to investigate the biological function  
24    of the blue pigment. Genetic analyses based on the mapping of the random insertions allowed the  
25    identification of eight genes involved in pigment production, including the second copy of *trpB*  
26    (*trpB\_1*) gene. Phenotypic characterisation of Ps\_77 white mutants demonstrated that the blue pigment  
27    increases oxidative-stress resistance. Indeed, while Ps\_77 was growing at a normal rate in presence of  
28    5 mM of H<sub>2</sub>O<sub>2</sub>, white mutants were completely inhibited. The antioxidative protection is not available  
29    also for non-producing bacteria in co-culture with Ps\_77.

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## 31    **Keywords:**

32    *Pseudomonas fluorescens*, transposon mutagenesis, blue-pigment biosynthesis, oxidative stress  
33    resistance, tryptophan metabolism.

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## 1. Introduction

*Pseudomonas fluorescens* is a well-known food spoiler and, although it is not commonly considered a human pathogen, it is an important species in public hygiene and food industry (Scales et al., 2014). The interest in food spoiling *P. fluorescens* has increased since 2010, when some packages of mozzarella were seized after a blue discolouration was noticed on the external surface of the cheese (RASFF, Annual Report 2010). The strains responsible for the discolouration were identified as belonging to a specific genetic cluster of the *P. fluorescens* group (Andreani et al., 2014). This cluster, called the “blue branch”, was demonstrated to contain all the blue-pigmenting strains as well as some unpigmented strains. Recent characterization of other strains producing the blue discoloration confirmed their monophyletic origin (Chierici et al., 2016).

The chemical nature of the blue pigment has been investigated, but no clear answer has been obtained yet. Based on a MALDI-TOF Mass spectrometry analysis, Andreani and colleagues (2015) suggested that the blue pigment may be an indigo-derivative. This result was also supported by a genomic investigation that revealed the exclusive presence of homologs of genes involved in tryptophan production in the genome of “blue branch” strains (Andreani et al., 2015); indeed, tryptophan has been reported to be involved in indole and indigo production in other bacterial species (Berry et al., 2002). The second copies of *trp* genes are included in a cluster of sixteen genes exclusive to the blue strains, identified in contig 4 of the genome of Ps\_77 and thus called c4\_BAR (Contig 4 Blue Accessory Region; Andreani et al., 2015).

The complete biosynthetic pathway, as well as the biological function of pigment production remain an open question. Several bacterial pigments, including *Pseudomonas* pigments, have a function as siderophore (Cornelis, 2010). Indeed, a previous study suggested a role for the blue pigment in iron metabolism (Andreani et al., 2015). However, some bacterial pigments have a role in oxidative stress resistance: for example, melanin produced by *Pseudomonas aeruginosa* confers resistance to oxidative stress (Rodríguez-Rojas et al., 2009) and this was also reported for the indigo pigment produced by

60 *Pseudomonas* sp. HAV-1 (Dua et al., 2014) and for the phenazine pigment in *Pseudomonas*  
61 *chlororaphis* GP72 (Xie et al., 2013).

62 In this study, we employ transposon mutagenesis (Goryshin et al., 2000) and characterize the white  
63 mutants to investigate the biosynthesis and the biological function of the blue pigment in strain Ps\_77.

64

## 65 **2. Materials and methods**

### 66 *2.1 Transposon mutagenesis and identification of Tn5-flanking sequences of selected mutants*

67 Detailed genomic and phenotypic information of Ps\_77 are already available (Andreani et al., 2015,  
68 LCYB000000000; SAMN03085510; GCF\_001542705.1). The wild-type strain and transposon-induced  
69 mutants were cultured and maintained in Luria Bertani Broth (LB; Sigma-Aldrich) or in Tryptic Soy  
70 Broth (TSB, Conda) and stored at -80 °C in LB Broth with 50% v/v glycerol (Sigma-Aldrich). Where  
71 appropriate, 50 ng/μL kanamycin sulfate (Sigma-Aldrich) was supplemented to the medium.

72 Transposon mutagenesis of Ps\_77 was carried out with EZ-Tn5™ Tnp Transposome™ (Epicentre, (an  
73 Illumina Company)®). Electrocompetent cells were prepared as reported by Choi and colleagues  
74 (2006). Cells were electroporated in an electroporation cuvette (ThermoScientific) with an Eporator  
75 (Eppendorf; 25 Uf, 200 Ω, 2.5 kV). Mutagenesis of Ps\_77 was performed twice to enhance the number  
76 of mutants losing the ability of producing the blue pigment. Screening of kanamycin sulfate resistant  
77 (Kan<sup>R</sup>) mutants was performed on Minimal Bacterial Medium Agar (MBM Agar, 7 g/L K<sub>2</sub>HPO<sub>4</sub>, 3 g/L  
78 KH<sub>2</sub>PO<sub>4</sub>, 0.5 g/L tri-sodium citrate, 0.1 g/L MgSO<sub>4</sub>, 1 g/L (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 2 g/L glucose and 15 g/L agar;  
79 Boles et al., 2004) with 50 ng/μL kanamycin sulfate as on this medium the production of the dark blue  
80 pigment is more evident than in complete media. Kan<sup>R</sup> mutants were picked based on the pigmentation  
81 and streaked three times on MBM Agar with 50 ng/μL kanamycin sulfate at 28 °C to check the  
82 phenotype. As a reaction to Kovac's reagent (isoamyl alcohol, para-Dimethylaminobenzaldehyde,  
83 concentrated hydrochloric acid; Sigma-Aldrich) was reported for blue pigmenting strains (Andreani et

84 al., 2015), phenotype was checked by striking suspected white colonies on one drop of this reagent.  
85 All selected Kan<sup>R</sup> mutants were grown in 5 mL of LB Broth with 50 ng/μL kanamycin sulfate and  
86 incubated overnight at 28 °C. Genomic DNA of selected strains was extracted using FastDNA™ SPIN  
87 Kit (MP BIOMEDICALS) and sequenced as described in Karlyshev et al., 2000 and Walterson et al.,  
88 2014. The PCR products were sent to MacroGen Inc. (Amsterdam, the Netherlands) for direct Sanger  
89 sequencing with the upstream primer (KAN-2 FP-3) or the downstream primer (KAN-2 RP-3). All the  
90 sequences were checked for quality and edited with FinchTV 1.4.0 software (Geospiza). The sequences  
91 of the primers are reported in Table 1S.

## 92 *2.2 Bioinformatic analyses of disrupted genes*

93 Insertion sequences were queried against NCBI using the Basic Local Alignment Search Tool  
94 (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>). HMMER was applied with the aim to highlight protein  
95 homology of the query against Reference proteomes or SwissProt (Finn et al., 2015). SIM (Local  
96 Similarity Program; Huang and Miller, 1991) was applied to align amino acidic sequences.  
97 Additionally, sequences were queried against Ps\_77 (LCYB000000000) through the Bioedit Blast Tool  
98 using the draft genome as database (Tippmann, 2004). Identification of COG (Cluster of Orthologous)  
99 was performed using KOALA (KEGG Orthology and Links Annotation; Kanehisa et al., 2015).  
100 Expression of the genes identified with the mutagenesis was evaluated using transcriptomic data  
101 obtained in a previous work (Andreani et al., 2015; SRR1725678; SRR1725679; SRR1725680;  
102 SRR1725681; SRR1725682; SRR1725683; SRR1725684; SRR1725723). The orthologues of c4\_BAR  
103 proteins were identified using STRING v10 (<http://string-db.org/>, Szklarczyk et al., 2015) and the  
104 reconstruction of the genomic region in each species was performed using Biocyc (<https://biocyc.org>).

## 105 *2.3 Evaluation of the transcription units of the contig\_4 blue accessory region*

106 c4\_BAR contains sixteen genes, of which the first fourteen, including *trp* genes, have the same  
107 orientation. The evaluation of the transcription units within the fourteen genes of the c4\_BAR was  
108 evaluated through PCR analysis. Thirteen primer pairs were designed using as template the sequence  
109 of Ps\_77 (LCYB000000000) between 3' and 5' of each couple of consecutive genes. Primers are

110 reported in Table 1S. RNA was extracted and retrotranscribed as described in Andreani et al., 2015.  
111 PCR amplifications were performed in an Applied Biosystems 2720 Thermal Cycler in a final volume  
112 of 20  $\mu$ L of amplification mix containing 1U of GoTaq polymerase (Promega, Madison, WI), 1X  
113 GoTaq Buffer, 1.5 mM  $MgCl_2$ , 0.2 mM each deoxynucleotide triphosphate (dNTP), 250 mM each  
114 primer and 1  $\mu$ L of cDNA as template. The PCR cycle was 94°C for 5 min and 35 cycles each 94°C  
115 for 30 s, 58°C for 30 s and extension at 72°C for 30 s. Amplified products were analysed by  
116 electrophoresis on 1.8% agarose-Tris-acetate-EDTA (TAE) gels, stained with SYBR® Safe DNA Gel  
117 Stain (Invitrogen, Carlsbad, CA) and visualized on a UV transilluminator (Gel Doc XRTM, Biorad).

#### 118 *2.4 Growth assays*

119 With the aim to investigate the biological function of the blue pigment, the phenotypic characterisation  
120 of Ps\_77 and its transposon insertion mutants was performed. All strains were grown overnight in LB  
121 broth at 28 °C in continuous shaking to reach an approximate count of  $10^8$  CFU/mL. Cultures were  
122 diluted to  $10^3$  CFU/mL in the final medium used for the assay to remove traces of LB broth for all the  
123 following phenotypic tests. All the tests were performed in triplicate. Growth curves were obtained  
124 using a plate reader (Multiskan Series Microplate Readers, Thermo Fisher Scientific), incubating the  
125 strains in MBM in a 96-well plate in continuous shaking at 22 °C and reading the optical density (O.D.)  
126 at 600 nm. Growth rate was calculated using DMFit tool of ComBase, an online tool allowing the  
127 shape of microbial growth curve, based on the Gompertz and Baranyi modified equation (Baranyi and  
128 Tamplin, 2004; Mytilinaios et al., 2012). Even if the model was initially created for the investigation  
129 of growth data in log CFU format, recent investigations revealed its applicability also to optical  
130 density-based growth curve (Mytilinaios et al., 2012; Rickett et al., 2015).

131 To assay whether the wild-type Ps\_77 and ~~the~~ four white mutant strains were able to use different  
132 carbon sources, a Biolog GN2 Micro Plate was used for each strain. Strains were grown overnight at  
133 22 °C in LB broth and a 5-fold dilution was pipetted in each of the 96-wells of the Biolog GN2 plate.  
134 Biolog GN2 Micro Plates were incubated at 22 °C and bacterial growth was evaluated after 24 hours

135 by measuring the O.D. at 600 nm with a plate reader. Growth of the wild-type and the selected mutants  
136 was evaluated as difference of the OD 600 nm of the strains in the control well (water).  
137 Growth assays were carried out in King's medium B broth (KB; 20 g/L of proteose peptone, Difco, 1,5  
138 g/L of K<sub>2</sub>HPO<sub>4</sub>, 1,5 g/L of MgSO<sub>4</sub>·7H<sub>2</sub>O, 10 mL/L of glycerol) with 100 µg/mL human apotransferrin  
139 (Sigma), a natural iron chelator, and 20 mM sodium bicarbonate, necessary for iron chelation by  
140 apotransferrin (Inglis et al., 2012) to obtain an iron-free environment. The production of the  
141 siderophore pyoverdine by the mutants was investigate after growth on MBM medium plates  
142 visualizing the fluorescence with a UV lamp (365 nm).  
143 To evaluate stress resistance, the growth of clones was measured in MBM at several concentration of  
144 hydrogen peroxide (H<sub>2</sub>O<sub>2</sub> 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, and 5.0 mM), NaCl (2.5, 3.0, 3.5, 4.0, 4.5,  
145 5.0, 5.5, 6.0, 6.5, 7.0 and 7.5%), different pH values (2.0, 2.5, 3.0, 3.5, 4.0 and 4.5) and after incubation  
146 of 2 hours at 55 °C and 4 °C. Growth curves were obtained using a plate reader (Multiskan Series  
147 Microplate Readers, Thermo Fisher Scientific), incubating the 96-well plate in continuous shaking at  
148 22 °C and reading the O.D. at 600 nm. The effect of hydrogen peroxide was evaluated also in a  
149 complete medium (TSB).  
150 All the statistical analysis was performed using Prism Graphpad software. The non-parametric  
151 Wilcoxon test was applied to compare the growth rates of the strains in all the tested conditions. The  
152 paired t-test was applied to compare the growth after 24 hours.

### 153 *2.5 Competition experiment*

154 Ps\_77 and M3 were grown separately in TSB until an OD<sub>600nm</sub> of 0.2 and then diluted 20-fold and  
155 mixed together (in a 1:1 ratio) in MBM at 2.5 and 5 mM concentration of H<sub>2</sub>O<sub>2</sub>. Competitions were  
156 set up in a 96-well microplate and incubated in continuous shaking at 22 °C. 100 µL of co-cultures 10-  
157 fold diluted in MBM broth were plated in MBM agar after 24 hours of incubation. Blue and white

colonies were counted after 4 days of incubation at 22 °C and paired t-test was applied to compare the results.

### 3. Results

#### 3.1 Generation and genetic characterization of white Ps\_77 mutants

Using the EZ-Tn5™ Tnp Transposome™,  $1.35 \cdot 10^5$  single individual transposon insertion mutants were obtained from the two different electroporation events. The combined application of MBM Agar and Kovac's reagent allowed an easy discrimination of the phenotype. White transposon mutants were picked and re-assayed on MBM agar to confirm stable insertion of Tn-5 transposon in the genome of Ps\_77. Ten transposon mutants showing a stable white phenotype were selected for further characterisation.

The genomic region flanking the transposon insertion sites was sequenced for each selected strain and the complete list of gene insertions observed in the investigated mutants is reported in Table 1. The localization of transposon-insertions allowed the identification of genes putatively involved in blue-pigment biosynthesis. Among the ten non-pigmenting mutants, insertions were localized in eight different genes. BlastKOALA tool allowed the annotation of only four out of eight genes, as reported in Table 1. The expression of these eight genes was checked on transcriptomic data from a previous study (Andreani et al., 2015). All these genes were expressed in MBM broth at 22 °C.

Six mutants (M2, M3, M4, M6, M7 and M19) lost the blue pigment production due to the disruption of four genes (PFLuk1\_0667, PFLuk1\_0668, PFLuk1\_0673 and PFLuk1\_0674) located in c4\_BAR. PFLuk1\_00667 was annotated as Tryptophan synthase beta chain *trpB\_1*. The PFLuk1\_00668 locus was annotated as hypothetical protein and no characteristic domain was identified using Hmmer. PFLuk1\_00673 and PFLuk1\_00674 were annotated as UDP-2-acetamido-2-deoxy-3-oxo-D-



181 glucuronate aminotransferase (Table 1). Low similarity to other species was revealed by a Blast  
182 analysis. Both proteins seem to have a characteristic domain, namely a DegT/DnrJ/EryC1/StrS  
183 aminotransferase domain. Excluding the DegT/DnrJ/EryC1/StrS aminotransferase domain, the two  
184 amino acid sequences are quite different. PFLuk1\_00673 and PFLuk1\_00674 lengths differs by six  
185 amino acids and SIM analysis revealed a low similarity between the two proteins with 31.6% identity  
186 in 345 residues overlap and gap frequency of 2.0%.

187 Three sequences were located outside the c\_4 BAR region. These genes were categorized as belonging  
188 to *Metabolic pathways/Amino acid metabolism (hisG, speA and trpB\_1)*, two of these are also involved  
189 in *Biosynthesis of secondary metabolites (hisG and trpB\_1)*. One gene (*gacS*) was classified as  
190 *Environmental information processing*.

191 PFLuk1\_00503 was mapped during the investigation of M5 and encodes for an ATP-  
192 Phosphoribosyltransferase and its ortholog in *P. fluorescens* A506 is *hisG*, a gene coding a protein  
193 involved in the biosynthesis of amino acid histidine. Another gene involved in amino acid metabolism  
194 and in particular in polyamines biosynthesis (PFLuk1\_05234) was identified mapping the mutation of  
195 M12. This locus encodes for the biosynthetic arginine decarboxylase, responsible of the synthesis of  
196 agmatine from arginine, a precursor of putrescine (Moore and Boyle, 1990). Another mutant (M15)  
197 was characterised by disruption of gene encoded by PFLuk\_02002, namely a signal transduction  
198 histidine-protein kinase BarA, annotated as *gacS* in *Pseudomonas* species. *gacS* belongs to the  
199 GacS/GacA two-component system, involved in the signal/transduction pathway. Finally, M20 was  
200 characterised by gene disruption of PFLuk1\_05423, annotated as Hypothetical protein by Prokka.  
201 Hmmer analysis showed a low level of similarity of this protein with *tfoX\_C* encoding for a  
202 competence protein. No particular phenotype has been reported in literature as a result of gene deletion,  
203 apart from the loss of competence, or the ability to take up free DNA from the environment (Smeets  
204 et al., 2006).

### 205 3.2 Genetic and functional organisation of c\_4 BAR region

206 Most of the genes included in the c\_4 BAR region, with the exclusion of *trp* genes, are part of a  
207 conserved cluster, with a similar genetic organization observed in some species of Actinobacteria, as  
208 revealed by HMMER and STRING investigations. The structure of the cluster in the different species  
209 is illustrated in Figure 1. To define the transcription units of c4\_BAR genes in Ps\_77, the cDNA was  
210 amplified using primers located in all the fourteen genes and a unique large operon was identified.

### 211 *3.3 Growth and phenotypic characterization of Ps\_77 and white transposon mutants*

212 Strains M5, M12, M15 and M20 were excluded from the phenotypic evaluation as the involvement of  
213 the four genes in other important pathways could influence the interpretation of results. The focus was  
214 addressed on the four mutants of the c\_4 BAR genes, excluding M6 and M7 as analogous to M4 and  
215 M2, respectively.

216 To test whether the blue pigment could be related to iron metabolism homeostasis, growth rate was  
217 measured for the wild-type strain Ps\_77 and all four white mutants in MBM, KB broth and iron limited  
218 KB at 22 °C, as well as in BIOLOG GN2 plates. The wild type Ps\_77 and the white mutants had  
219 statistically indistinguishable growths in all the media tested ( $p > 0.05$  for all the pair comparisons,  
220 Table 2S, Figure 1S). The wild-type strain produced the blue pigment in the three condition tested and  
221 the pigment production was unequivocally visible after 48 hours of incubation.

222 The white mutant colonies appeared fluorescent as the wild-type strain at UV lamp exposure  
223 demonstrating that the lack of blue pigment does not influence pyoverdine production.

### 224 *3.4 Stress resistance of Ps\_77 and white mutants*

225 The capability of the blue pigment to confer resistance to different types of stress as osmotic (salt),  
226 oxidative, thermal and high/low pH was tested comparing the growth of wild-type and the four white  
227 mutants in presence of the stress agents (H<sub>2</sub>O<sub>2</sub>, NaCl, high/low pH and incubation at 55 °C-4 °C). The  
228 results indicated that all stressors produced the same effect on the wild-type as on the unpigmented  
229 mutants (Fig. 2S), except for oxidative stress (Fig 2). The growth of the four mutants in presence of

230 hydrogen peroxide was slightly reduced at 2.5 mM of H<sub>2</sub>O<sub>2</sub> and completely inhibited at 5 mM resulting  
231 at both concentrations significantly different from the growth of the wild-type (Wilcoxon test,  $p <$   
232 0.05). Figure 2 reports the lower ratio of growth of the four mutants *versus* the wild-type at two  
233 concentrations (2.5 and 5 mM) of hydrogen peroxide after 24 hours of incubation. Similar effect of  
234 resistance to hydrogen peroxide by the blue wild-type strain occurred in complete medium (TSB,  
235 Figure 2A) and minimal medium (MBM, Figure 2B).

236 As blue pigment is released extra-cellularly, a co-culture of Ps\_77 and M3 strains was set up in MBM  
237 liquid medium added with hydrogen peroxide at 2.5 and 5 mM to evaluate whether pigment mediating  
238 antioxidative protection have the ability to protect non-producing bacteria. The growth of the white  
239 mutant was significantly lower than the wild type (paired t-test  $p < 0.05$ , Figure 3) indicating a not  
240 diffusible antioxidant effect of the blue pigment in the culture.

#### 241 **4. Discussion**

242 The present work aimed to investigate the biosynthetic pathway of the blue pigment, as well as its  
243 biological function in one well-studied *P. fluorescens* strain through the creation of a library of non-  
244 pigmenting strains and their genotypic and phenotypic characterization. A high number of mutants of  
245 *P. fluorescens* Ps\_77 was created through the application of EZ-Tn5™ Tnp Transposome™ kit.  
246 However, only ten showed a stable insertion, as showed by the phenotype observed on MBM agar and  
247 after testing with Kovac's reagent. The negative reaction of Kovac's reagent of the white mutants  
248 confirms that the blue pigment or its precursor is an indole-derivative. It has been in fact demonstrated  
249 that this reagent can produce different coloured substances when reacting with different indole  
250 derivatives (Ehmann, 1977).

251 The transposon mutagenesis identified eight genes involved in blue pigment production in Ps\_77 and  
252 among these four are included in c\_4 BAR region. This result confirms previous hypothesis that  
253 indicated the involvement of the second copies of *trp* genes in the blue pigment biosynthesis through

254 an indole/indigo biosynthesis pathway (Andreani et al., 2015). Carriage of multiple *trpB* gene copies  
 255 has been reported in several bacterial and archaeal species (Xie et al., 2001; Merkl, 2007; Busch et al.,  
 256 2014) and are frequently involved in biosynthesis of molecules with different functions, despite the  
 257 high-energy cost required for this pathway. It has been demonstrated that usually the second copy  
 258 possesses substrate specificity, not assembled with TrpA and seems to have other functions, such as  
 259 indole salvage (Busch et al., 2014; Hiyama et al., 2014). In *Pseudomonas aeruginosa* the first enzyme  
 260 of the tryptophan biosynthesis pathway (Antranilate synthase coded by *trpE* and *trpG* genes) is  
 261 duplicated and the second copy is involved in the quorum sensing signalling but not in tryptophan  
 262 biosynthesis (Palmer et al., 2013). Moreover, indole and the indole-derivatives, synthesized from  
 263 tryptophan are interspecies and interkingdom signalling molecules involved in important roles as  
 264 bacterial pathogenesis and eukaryotic immunity (Lee et al., 2015).

265 Only a *trpB\_1* mutation was identified as conferring a white phenotype in Ps\_77. The second copies  
 266 of other *trp* genes (*trpA*, *trpC*, *trpD*, *trpF*) were not identified to be disrupted in the white mutants.  
 267 This result can have several explanations. The mutation in the other *trp* genes might be complemented  
 268 by the first copy of the aforementioned genes or only TrpB might be the only one directly involved in  
 269 the pigment production. A third possible explanation might be that the mutants isolated were not  
 270 enough to identify all the genes involved in pigment production. The other three white mutants with  
 271 the disrupted gene located c\_4 BAR are a hypothetical protein (PFLuk1\_0668) and two  
 272 aminotransferases (PFLuk1\_0673 and PFLuk1\_0674). DegT/DnrJ/EryC1/StrS aminotransferases  
 273 have been widely reported as having a regulatory and protein kinase (sensor) function (Murphy et al.,  
 274 1993; Madduri and Hutchinson, 1995). In *Bacillus stearothermophilus*, *degT* has a proper regulatory  
 275 function, being involved in the transfer of environmental stimuli (Takagi et al., 1990).  
 276 Aminotransferases of this family have been identified as involved in aminotransfers that lead to amino  
 277 sugars involved in the formation of LPS and aminoglycosides in *Porphyromonas gingivalis* (Shoji et  
 278 al., 2002). On the other hand, Chen and colleagues (2000) speculate that the lower blue pigment

279 production in *P. gingivalis degT* mutants might be due to the impossibility of the mutant strain to  
280 secrete the extracellular pigment through the production of vesicles. Excluding the  
281 DegT/DnrJ/EryC1/StrS domain, the two proteins (PFLuk1\_0673 and PFLuk1\_0674) are quite  
282 different. This result induces to suppose the two loci have different functions, even if sharing the same  
283 domain and might be involved in the secretion of the blue pigment or of a precursor of the blue  
284 pigment.

285 In species belonging to three different families of Actinobacteria (Streptomicetaceae,  
286 Pseudonocardiaceae and Nocardopsaceae), the three genes PFLuk1\_0668, PFLuk1\_0673 and  
287 PFLuk1\_0674 are included, with their neighbours, in c\_4 BAR (excluding *trp* genes), in a large gene  
288 cluster. The function of these proteins is unknown. Several of these Actinobacteria species were shown  
289 to be producers of a rich array of active metabolites (Li et al., 2013; Girard et al., 2014). The c\_4 BAR  
290 identified in Ps\_77 comprised the nine Actinobacteria homologue genes assembled with the five *trp*  
291 genes. The analysis of the transcripts in Ps\_77 demonstrates that all the fourteen genes are co-expressed  
292 and are included in a single large operon supporting the hypothesis that these genes are coding the  
293 enzymes for the biosynthesis of the pigment and related function as transport and secretion.

294 Among the mutants disrupted in genes located outside c\_4 BAR, two are involved in amino acid  
295 metabolism (*hisG* and *speA*). The effect of these mutations in pigment production it is not evident  
296 despite HisG and SpeA proteins are included in biosynthetic pathway that share intermediate  
297 compounds with *trp* metabolism. (i.e. Phosphoribosyl pyrophosphate; PRPP).

298 On the contrary, the implication of *gacS* product in blue pigment biosynthesis regulation it is not  
299 difficult to suppose. In fact GacS/GacA system has been reported to regulate several phenotypes in  
300 bacteria. In *Escherichia coli* and *Vibrio fischeri* the siderophore-mediated iron sequestration is  
301 regulated by GacS/GacA system (Sahu et al., 2003; Foxall et al., 2015). In *Pseudomonas fluorescens*  
302 FD6 (a biocontrol strain), it has been demonstrated that *gacS* knock-out strains cannot produce a wide

303 range of secondary metabolites. In particular, biofilm formation and siderophore production were  
304 downregulated, as well as the protection activity against *Botrytis cinerea* HY2-1 (Chang et al., 2014).  
305 To summarize, the blue pigment biosynthetic pathway is strictly related to *trp* genes, confirming the  
306 importance of the genomic region previously identified in c\_4 BAR of Ps\_77. For this reason, the  
307 phenotypic characterization of the mutants was focused on the four mutants with the disruption located  
308 in c\_4 BAR. Specifically, the study of the four mutants could better clarify the function of the pigment  
309 better than the other mutants as *GacS*-, *hisG*- and *speA*-knock-out strains that are involved in other  
310 important pathways.

311 Wild type and mutant strains grow with a comparable growth rate in normal medium (MBM; KB),  
312 iron-depleted medium and different carbon sources. This suggests that the lacking of the pigment does  
313 not affect the growth capability or the utilisation of different carbon sources as well as the production  
314 of the pigment is not induced by the absence of iron in the medium, suggesting that the pigment is not  
315 a siderophore (Cornelis, 2010).

316 Several natural pigments produced by bacteria, fungi or microalgae have stress resistance properties  
317 (Tuli et al., 2015). No effect on osmotic, thermic and pH stress-resistance was recorded. However,  
318 Ps\_77 demonstrated an increased resistance to oxidative stress induced by hydrogen peroxide. This  
319 result suggests that the blue pigment (or its parent compound) might act as an antioxidant agent or  
320 regulates an antioxidant response and is not involved in a general stress response. Moreover, the  
321 involvement of GacA/GacS system in oxidative resistance in *P. fluorescens* (Heeb et al., 2005) and the  
322 antioxidant activity of indole, indole derivatives and indigo (Dua et al., 2014, Lee et al., 2015) support  
323 this hypothesis.

324 The capability of surviving to oxidative stress is of great importance to survive in the environment.  
325 Several strains of *Pseudomonas fluorescens* are component of the rhizosphere, an environment in  
326 which the active metabolism of the root tip generates a high amount of reactive oxygen species. A  
327 study demonstrated that the success of root colonization in *P. putida* depends on its capability to resist

328 to oxidative stress (Kim et al., 2000). Bacterial tryptophan metabolites interfere with immune  
329 responses in plants and animals and kynurenine pathway may allow immunomodulatory interplay  
330 between bacteria and host (Genestet et al., 2014, Lee et al., 2015, Bortolotti et al., 2016).

331 The competition experiment demonstrated that the resistance to hydrogen peroxide is limited to the  
332 blue producing strain and it cannot be shared with strains sharing the same environment. The diffusion  
333 of the blue pigment in the plate or liquid medium occurs despite their insolubility in water (Andreani  
334 et al., 2015). This means that the diffusion of the blue pigment in the medium has to be supposed  
335 complexed with other compounds. For this reason, it would be likely to assume that the blue pigment  
336 in this form cannot act as antioxidant agent. The antioxidant activity might be exerted by the blue  
337 pigment or its precursor inside the bacterial cell or linked to the plasmatic membrane.

## 338 **5. Conclusions**

339 The data obtained in the present study strongly suggest a role in resistance to oxidative stress of the  
340 blue pigment, despite its chemical structure was not completely elucidated. The pigment, that was  
341 confirmed to be an indole derivative, is produced and secreted in the environment. This capability  
342 successfully adapts the blue-producing strains to survive in the different environment from which *P.*  
343 *fluorescens* is frequently isolated as polluted environment, rhizosphere or frequently sanitized food  
344 industries. The elimination of blue producing strains of *P. fluorescens* from the production line remains  
345 an unresolved problem for dairy industries. The increased resistance to antioxidant agents might help  
346 explain the difficulty of eradication.

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499 **Captions**

500 **Figure 1.** Gene cluster structure of the c\_4 BAR gene homologues in Actinobacteria: *S.*  
501 *griseoaurantiacus* M045 (*Streptomycetaceae*); *Streptomyces* e14 (*Streptomycetaceae*); *K. setae*  
502 KM6054 (*Streptomycetaceae*); *A. mirum* DSM43827 (*Pseudonocardiaceae*); *S. erythraea* NRRL2338  
503 (*Pseudonocardiaceae*); *N. dassonvillei* DSM43111 (*Nocardiopsaceae*). Above each gene, the  
504 locus\_tag is reported.

505 **Figure 2.** Ratio of the growth of mutants (M2, M3, M4, M19) and Ps\_77 in TSB (A) and MBM (B)  
506 added with H<sub>2</sub>O<sub>2</sub> at 2.5 and 5 mM after 24 hours (\* p < 0.05).

507 **Figure 3.** Ratio of the bacterial cell counts of M3 mutant and Ps\_77 in MBM added with H<sub>2</sub>O<sub>2</sub> at 2.5  
508 and 5 mM (\* p < 0.05).